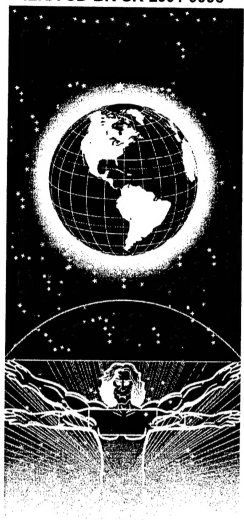
IERA-SD-BR-SR-2001-0006



UNITED STATES AIR FORCE IERA

The Influence of Heterogeneity
in Gamma Spectroscopy Analysis
of Soil Contaminated with Weapons
Grade Plutonium at the BOMARC
Missile Accident Site,
McGuire AFB NJ

Steven E. Rademacher, Major, USAF, BSC

20020108 122

October 2001

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Air Force Institute for Environment, Safety and Occupational Health Risk Analysis Surveillance Directorate Radiation Surveillance Division 2513 Kennedy Circle Brooks Air Force Base TX 78235-5116

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REPORT DOCUMENTATION PAGE

Form Approved OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Washington headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Pagetryork Reduction Project (0704-0188), Washington, DC 20503.

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507 SYMINGTON DRIVE											
SCOTT AFB IL 62225-5022		IE	RA-SD-BR-SR-2001-0006								
11. SUPPLEMENTARY NOTES											
12a. DISTRIBUTION AVAILABILITY		12b. DI	STRIBUTION CODE								
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List of Acronyms

AFIERA - Air Force Institute for Environment, Safety and Occupational Health Risk Analysis

BOMARC - Boeing Michigan Aeronautical Research Center

CI - confidence interval

EPA - Environmental Protection Agency

FIDLER - field instrument for detecting low energy radiation

HpGe - hyperpure germanium

HQ AFSC - Headquarters, Air Force Safety Center

HQ AMC - Headquarters, Air Mobility Command

NaI(Tl) - thallium-drifted sodium iodide

LANL - Los Alamos National Laboratory

LLL - Lawrence Livermore Laboratory

MDC - minimal detectable concentration

NAEG - Nevada Applied Ecology Group

NIST - National Institute for Standards and Technology

NTS - Nevada Test Site

ORNL - Oak Ridge National Laboratory

% CV - percent coefficient of variation

REECo - Reynolds Electrical and Engineering Company

RPD - relative percent difference

WGP - weapons grade plutonium

Abbreviations

Am - americium
cm - centimeter
°C - degree Centigrade
g - gram
h - hour
keV - kilo electron volt
μm - micrometer
mm - millimeter
MeV - million electron volt
$\boldsymbol{\mu}$ - linear attenuation coefficient
n - number
pCi - picocurie
Pu - plutonium
ρ - density
R ² - squared correlation coefficient
σ - standard deviation
s - second

y – year

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Introduction

The Boeing Michigan Aeronautical Research Center (BOMARC) Missile Site is an inactive Air Force installation located in Plumstead Township, New Jersey. The site was an active nuclear missile defense site from 1958 – 1971. On June 7, 1960, a fire occurred in one of the shelters in which the shelter, missile, and warhead were partially consumed by the fire. The high explosive materials in the weapon ignited but did not detonate. The most intense period of the fire lasted about one hour. Water was applied to the shelter and weapons during the fire by the installation fire department. The fire melted the weapons grade plutonium (WGP) that was contained in the device. Turbulent local atmospheric conditions and the water applied during the fire contributed to scattering of WGP to the environment. WGP is the primary radiological concern at the site. The amount of WGP originally contained in the weapons remains classified. Los Alamos National Laboratory estimated that approximately 100 g of WGP remained on-site (Farley 1996). A recent characterization survey supports an estimate closer to 300 g (OHM 1998).

Quantification of WGP in soils is a technically challenging endeavor compared to many naturally-occurring or other man-introduced radiological contaminants. A large source of the difficulty with sampling and analysis of soil samples is related to the discrete particulate nature of plutonium contamination (referred to as "heterogeneity" in this report) (Bernhardt 1976). In the 1960's and early 1970's, the preferred analytical method for soil samples containing WGP was chemical dissolution and α -spectroscopy analysis. Because of practical limitations in aliquot masses prepared for dissolution (i.e., less than 50 g), the method was plagued with large uncertainties if plutonium contaminants were in discrete particle form. The uncertainties from this method were largely due to the inability to collect samples representative of the contaminated area rather than from systemic or random laboratory errors.

With improvements in detector technologies, alternate methodologies were developed. The Field Instrument for Detecting Low Energy Radiation (FIDLER) is a specialized NaI(Tl) detector designed for in-situ measurements of WGP (Tinney 1969). Later, high-resolution γ-spectroscopy systems were commonly used for laboratory assessment of plutonium activity concentrations from ²⁴¹Am quantification and knowledge of the relationship of the ²⁴¹Am and plutonium. This method, while dependent on accurate assessment or knowledge of the plutonium to ²⁴¹Am activity concentrations, can be used to evaluate samples with mass up to a kilogram. The ability to evaluate sample masses of this magnitude offers significant advantage over chemical dissolution and α -spectroscopy analysis where sample mass is highly limited. First, the γ -spectroscopy method does not require chemical procedures that are labor intensive and time consuming. Second, the larger sample mass ensures that the sample is more representative of the environmental area being evaluated. On a negative note, however, little research has been conducted to investigate the effect of heterogeneity on uncertainties on laboratory γ-spectroscopy analysis assessments. This report summarizes the results of a sampling and laboratory analysis effort on BOMARC site soils. The primary purpose of the study was to better understand the effects of heterogeneity on sample collection and analysis through y-spectroscopy of a WGP contaminant. The study demonstrated that the particulate nature of the WGP contaminant created significant levels of uncertainty in quantified activity concentrations for some samples evaluated. The impact of the uncertainties is largely dependent on the purpose of the assessment. For a site characterization

effort, a high degree of accuracy in analytical results may not be necessary. In this case, the impact is negligible. For a research study, however, uncertainties from a heterogeneously distributed contaminant may greatly limit the purpose of a study. For site closure measurements, where post remediation residual WGP contamination may be very low, uncertainties may be overwhelming and greatly limit a conclusion on the effectiveness of the effort. In addition, for low activity concentration samples, overall uncertainty will be exacerbated by that introduced by counting uncertainties alone. A method to reduce the effects of heterogeneity on sample analysis is provided.

Background

1. Weapons Grade Plutonium.

WGP is comprised primarily of ²³⁹Pu, with lesser mass amounts of ²³⁸Pu, ²⁴⁰Pu, ²⁴¹Pu, and ²⁴²Pu. For many nuclear weapons, individual WGP isotopic assay information exists (classified). However, for the warhead on the BOMARC missile and other nuclear weapons produced during the same time, specific assay information is not available (Taschner 1998). Table 1 provides an estimate of the isotopic composition of the BOMARC missile based on information from Los Alamos National Laboratory (Taschner 1998) and soil analyses performed in 1997 (Rademacher 1999).

Table 1. Isotopic Composition (by mass) of WGP in BOMARC Weapon Based on Los Alamos National Laboratory Estimates and Soil Analyses (Rademacher 1999).

Isotope	Mass Fraction*	Radiological Half-life (y) **						
Pu-238	0.0099	87.74						
Pu-239	0.937	24,110						
Pu-240	0.056	6,560						
Pu-241	0.0047	14.35						
Pu-242	Negligible	376,000						

^{*} Fractions in 1958 ** Walker et al 1984

There was variability in the isotopic composition of WGP among weapons produced during this period because of variables involved in WGP production. According to Los Alamos National Laboratory records (Taschner 1998), the plutonium in the BOMARC missile (involved in the fire at McGuire AFB) may have been separated in 1957/1958. The relative isotopic composition of WGP constituents changes over time due to radioactive decay. Shortly after chemical separation during production, the most significant change is due to the radioactive decay of ²⁴¹Pu:

241
Pu $\rightarrow ^{241}$ Am $+ {}_{-1}^{0}\beta$.

The daughter product, 241 Am, is an α -particle emitter with a radiological half-life of 432 y (Walker et al 1984).

Table 2 lists the major radiation(s) emitted by the primary constituents of WGP. For ²³⁹Pu and ²⁴⁰Pu, only infrequent low-energy photons are emitted. Direct assessment of either of these isotopes in samples can be accomplished through high-resolution gamma spectroscopy if sample activities are reasonably high. Measurement accuracy can be severely affected if the density and elemental

composition of the sample are not well characterized. Better measurement accuracy can be obtained through chemical dissolution, separation, and alpha or mass spectroscopy. However, this measurement approach has shortcomings. To ensure measurement accuracy, the sample must be completely dissolved by strong acids. The requirement for complete dissolution is time consuming and limits sample mass. Many laboratories limit sample mass to about 10 g, with many commercial environmental laboratories using less than 2 g. Further, application of acid leaching methods on BOMARC soils was effective in solubilizing only about 50 % of the contaminant (Refosco 2001). Small sample mass can introduce variability into the activity assessment if WGP is heterogeneously distributed. Most importantly, the method is expensive because it involves wet chemistry and can be cost-prohibitive for a large number of samples. Because the α -particle energies of the 239 Pu and 240 Pu are very close, α -spectroscopy is incapable of resolving the two isotopes. Practically, for radiation protection purposes this does not present a problem because each isotope has the same activity to dose conversion factor (Eckerman 1999).

Table 2. Major Radiation Emissions of WGP Constituents (Scheien 1992).

	α-Particle Energies	β-Particle Energies	Photon Energies				
Radionuclide	(MeV) & Frequency	(MeV) & Frequency	(MeV) & Frequency				
	5.155 (0.733)		0.113 (0.0005)				
Pu-239	5.143 (0.151)	None	0.014 (0.044)				
	5.105 (0.115)						
Pu-240	5.168 (0.735)	None	0.054 (0.0005)				
ru-240	5.123 (0.264)	None	0.014 (0.11)				
Pu-241	None	0.021 (1.00)	None				
	5.486 (0.852)		0.014 (0.427)				
Am-241	5.443 (0.128)	None	0.0595 (0.359)				
	5.388 (0.014)		0.026 (0.024)				

Of the radionuclides in WGP, 241 Am has the most favorable photon emission characteristics with a 0.0595 MeV γ -ray at an emission frequency of 36 %. This photon is of sufficient energy to afford reasonable measurement sensitivity in high-resolution gamma spectrometry counting systems. Additionally, the attenuation is not as severe as that of the 17 keV x-ray emitted by 239 Pu or 240 Pu. In laboratory and in-situ environmental measurements of WGP, measurement of the 0.0595 MeV γ -ray from 241 Am is often used as a surrogate for $^{239/240}$ Pu, provided the ratio of activities between the two elements is well known.

2. Plutonium Characteristics in the Environment.

Plutonium is a silvery-white metal that readily oxidizes in air. The compound can combine with oxygen to form many different compounds to include binary oxides, peroxides, hydroxides, and oxides of higher order. Plutonium dioxide, PuO₂, is the most stable of the oxides found in the environment and is formed under most conditions, especially when plutonium metal is ignited in air (Burley 1990). PuO₂ has a high melting point (2240 °C), has a high chemical stability, and is highly

insoluble in water (Burley 1990). The behavior of plutonium in soils can vary depending on the local soil characteristics and the form the plutonium is in at the time of introduction. Four sites have been extensively monitored in the U.S. up to 30 years after introduction of the plutonium into the environment: the Nevada Test Site (NTS), Oak Ridge National Laboratory (ORNL) in Tennessee. Mound Laboratory in Ohio, and Rocky Flats in Colorado (Burley 1990). The source of plutonium is different for each site. At NTS, the plutonium is dispersed as an oxide as the result of safety research studies. At Oak Ridge, plutonium in a holdup pond was released when a dike broke. At Rocky Flats, cutting oil contaminated with metallic plutonium was released from leaking storage drums; while at the Mound Facility, a low-pH solution of plutonium leaked from a waste transfer line. For the soils studied at NTS and Rocky Flats, extraction of plutonium from soils was very low (10-15%) as compared to 60-85% extraction from the soils of the Mound Facility and ORNL (Burley 1990). Thus, if the plutonium was introduced into the environment as an oxide or metallic form, it exhibited low solubility; whereas if introduction was in the form of a soluble compound. much greater mobility was exhibited at a later time. Furthermore, autoradiographic comparisons of Rocky Flats and Mound Facility soils have indicated that the soils of the former exhibited discrete particles of plutonium, while the latter had a more homogeneous dispersion (Burley 1990). Homogeneous dispersion of plutonium requires the material to be available in an ionic form. For plutonium at the Mound Facility, the release was in the form of an acid solution, which allowed a more uniform dispersal of the plutonium into the soil matrix. Because of the nature of the release. the plutonium at the BOMARC site is expected to exhibit characteristics similar to that released at Rocky Flats and NTS.

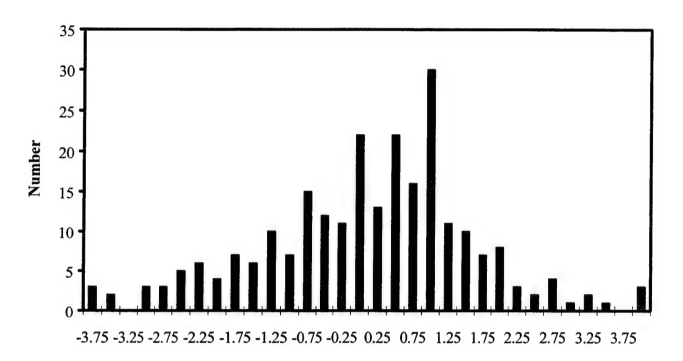
3. Studies of Plutonium at the BOMARC Site.

There has been a considerable number of environmental surveillance activities on the BOMARC site over the past 40 years. AF personnel accomplished a majority of the surveillance, but in the last 10 years most of the activities have been accomplished by organizations under contract to the AF. Three recent reports on surveillance activities note problems associated with evaluation of soil concentrations because of heterogeneity effects (HQ AMC 1992, Watts and Collins 1992, Kennedy 1990). The first two references did not provide data to support problems associated with heterogeneity. The last reference provided actual site data. The data was comprised of in-situ γ -measurements with FIDLER and analysis for ²⁴¹Am using laboratory γ -spectroscopy with a hyperpure germanium (HpGe) detection system. The purpose of the study was to determine a relationship between the FIDLER and ²⁴¹Am activity concentrations. A regression of the two parameters provided a squared correlation coefficient of 0.171. The author concluded, "it appears that there is not a linear relationship between the data set." Likely, the lack of correlation is due to inability of the sampling and analysis methods to control the effects of heterogeneity.

In 1997, OHM Remediations Services Corp. performed a characterization effort to support the Remedial Investigation/Feasibility Study (RI/FS). The purpose of the characterization effort was to determine the extent of the contamination zone. Soil samples were analyzed by both α - and γ -spectroscopy. Many of the 249 samples had analyses for ²⁴¹Am accomplished through both methods. Figure 1 contains a histogram of the ratio of the ²⁴¹Am activity concentration for γ - to α -spectroscopy analyses. Due to the wide range of ratio values, the plot is of the logarithm of the ratios. The ratios ranged from 0.00049 to 2629, spanning more than six orders of magnitude. The

mean, median, and percent coefficient of variation (% CV) of the data are: 15.8, 1.17, and 1100 %, respectively. The evaluation of this data set was not provided in the characterization report. Headquarters, Air Force Safety Center (HQ AFSC) provided additional evaluations of the characterization study data (Rademacher 1999a). In general, the data evaluations from the HQ AFSC report noted high variability and the influence of heterogeneity.

Figure 1. Logarithm of Activity Concentration Ratio (γ - to α -Spectroscopy) for 241 Am (OHM 1998).



Log of Ratio (γ- to α-Spectroscopy) for Am-241

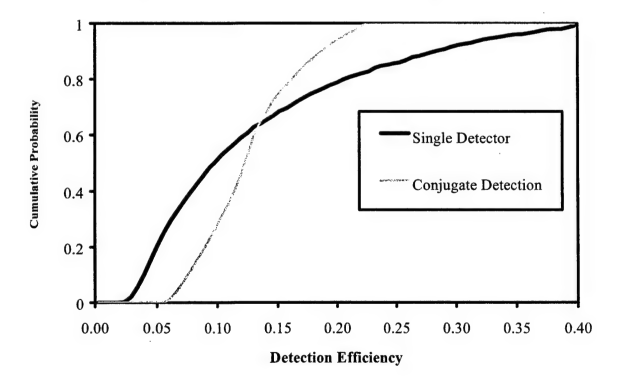
The high variability of the data in Figure 1 may be the result of many factors. First, HQ AFSC speculated that the α -spectroscopy evaluations were based on batch chemical recovery fractions in lieu of the preferred individual aliquot recovery method (Rademacher 2000a). The primary motivation for use of the former analytical method is reduced cost, because an ²⁴³Am tracer is used in a fraction of the aliquots rather than all. Heterogeneity is the other factor that is believed to have a significant influence on the variability exhibited in the data of Figure 1. Due to the particulate nature of WGP, and the aliquot masses used for α -spectroscopy (\sim 2 g) and γ -spectroscopy (\sim 500 g), this factor is likely the most important.

In 2001, the Nuclear Engineering Department of Massachusetts Institute of Technology investigated the chemical nature of two BOMARC site samples (Plaue and Czerwinski 2001). In general, the soils were highly resistant to chemical attack, suggesting that the majority of the plutonium and americium contaminants were in oxide forms. For the BOMARC environment, it was predicted that

about 15 % of the contaminants are in soluble carbonate phase, indicating that the BOMARC plutonium has similar characteristics to that of NTS plutonium.

4. Theoretical Predictions of the Effects of Single Particle Heterogeneity on γ-Spectroscopy Analysis of WGP. HO AFSC performed theoretical measurements to determine the effects of heterogeneity in WGP contamination in soils on y-spectroscopy analyses (Rademacher 2000b). The analysis was performed for varied soil volumes, linear attenuation coefficients (u), and sample containers. The calculations were made under the assumption that all of the WGP in a sample was contained in a single particle. This represents worst-case heterogeneity, while complete homogeneity represents the opposite. Figure 2 contains a plot of example detection efficiency calculations for a HpGe system and petri dish sample containers of similar dimension to those used in this study. The u for this set of efficiency calculations was 0.38 cm⁻¹, while the referenced report contains calculations for additional values. The solid black line of the plot is the cumulative frequency distribution of detection efficiency for a single HpGe detection system. From the data, detection efficiency ranges from 0.025 to 0.4, had a mean of 0.13, and % CV of 76 %. The solid gray line of the plot is the cumulative frequency distribution of the mean detection efficiency for conjugate measurements (i.e., counted on both sides). For this distribution, mean detection efficiency for the paired calculations ranged from 0.06 to 0.23, with a distribution mean of 0.13, and a % CV of 30 %. Clearly, conjugate counting provides significant reduction in variability for measurement of soil samples with maximum heterogeneity. In this example, variability is reduced by almost four-fold.

Figure 2. Cumulative Frequency Distribution of Detection Efficiency for a Single Particle in a 126 cm³ Volume Container ($\mu = 0.38$ cm⁻¹).



5. Variability in α-Spectroscopy Analysis of Soils with WGP Contaminants.

- a. <u>General</u>. While the primary purpose of this report is to better understand the effects of heterogeneity on γ -spectroscopy analysis of BOMARC soils, a summary of some observed effects on α -spectroscopy analyzed soils provides additional insight on the problems associated with heterogeneity. The analyses were of samples associated with investigation of WGP from NTS.
- b. Nevada Applied Ecology Group (NAEG). From the NAEG, conflicting analytical data from investigations was observed on samples being analyzed for WGP contaminants (Bernhardt 1976). To further investigate the problem, one sample was split among three laboratories: Environmental Protection Agency (EPA), Reynolds Electrical and Engineering Company (REECo), and Lawrence Livermore Laboratory (LLL). Each laboratory further blended and homogenized the sub-sample, split the sub-samples into smaller aliquots of varying mass dependent on the laboratory, chemically dissolved the sample, and analyzed the plutonium extract by α-spectroscopy. The results of the comparison are in Table 3. From the data in the table, it is apparent that significant variability existed among the reported mean activity concentrations of the laboratories, with a range of 1.1 to 5.3 pCi g⁻¹. Also, the study demonstrated that within laboratory variability was high and was generally inversely related to aliquot mass. For the EPA data, the percent coefficient of variation (% CV) was 130 %. Even for the LLL data, where aliquot masses were high, relative percent difference (RPD) values were high among some paired analyses.

Table 3. Comparative Analysis of ^{239/240}Pu in Soil from NTS (Adapted from Bernhardt 1976).

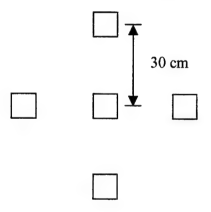
Lab	Aliquot Size	Number of	^{239/240} Pu	(pCi g ⁻¹)	%	^{239/240} Pu Range					
	(gram)	Aliquots	mean	σ	CV	(pCi g ⁻¹)					
EPA	1	14	1.1	1.4	130	0.23 - 5.3					
REECo	10	10	2.3	1.5	66	0.66 - 5.2					
	25	2	3.3	NA	83*	1.9 – 4.6					
LLL	25	2	3.4	NA	6*	3.3 - 3.5					
	100	2	4.8	NA	29*	4.1 - 5.5					
	100	2	5.3	NA	48*	4.0 - 6.5					

NA = Not Applicable * Relative Percent Difference

Methodology

- 1. Sample Collection. With a FIDLER instrument, the survey team scanned areas of the BOMARC site that did not contain concrete or asphalt overburdens. Areas with elevated instrument response were selected for sampling. Seven locations had five surface samples collected for a single FIDLER instrument measurement according to the spatial pattern of Figure 3. The surface samples were collected from the surface to a depth of 5 cm (2 in) with a stainless steel trowel. Between samples, the trowel was cleaned with distilled water. Samples were placed in plastic bags, with notation of sample number and depth. Sample chain-of-custody forms were prepared for the sample set. Prior to packaging for shipment, the samples were double-bagged. For the center locations in the sampling pattern, a sample was collected at a depth of 5-10 cm. For one center sample, a third sample at depth (10-15 cm) was collected. Twenty-three locations had only a surface soil sample collected in the center sub-sampling location. Ten locations had only a surface (0-5 cm) and a sample at depth (5-10 cm) that was collected in the center sub-sampling location. These 10 locations were sampled one year later than the other locations, with the intent of filling a data gap in low activity samples.
- 2. FIDLER Measurements. The FIDLER device used for the measurements had an energy window set to correspond to the 60 keV γ -ray from ²⁴¹Am. The detector was held in a stand, with a detector to ground surface separation distance of 30 cm. One-minute measurements were recorded.

Figure 3. Soil Sampling Pattern.



3. Sample Preparation. At the laboratory, the samples had mass determinations and filtration through a course mesh sieve to remove rocks. The remaining fraction was ashed in an oven at 500 °C for 24 h. The samples were blended and homogenized in a soil tumbler. The soil was incrementally placed into plastic petri dishes (9.5 cm in diameter x 2.54 cm in height) until completely filled, exhausting all of the soil. Due to variability in the mass of samples collected and rock fraction, aliquot number varied from 2 to 10 among each sample. Net sample mass was determined for each aliquot, with the typical being about 200 g. The petri dish lids were permanently sealed to the bases with a silicone sealant. The sample container exterior was cleaned and placed in a clean Zip-LockTM container to prevent contamination of laboratory surfaces.

- 4. Analysis. Aliquot 241 Am activity concentration was assessed by laboratory γ -spectroscopy analysis on a 8.9 cm diameter HpGe detector encased in a 7.5 cm thick steel cave. The aliquot to detector face separation distance was about 4 mm. Each aliquot was assessed twice, with the sample aliquot containers being flipped between measurements. Measurement periods were set at 10,000 s (2.8 h). Absolute activity concentration was based on comparison to a prepared soil standard that was calibrated against a National Institute of Standards and Technology (NIST) traceable calibration source.
- 5. Data Evaluation. Estimated aliquot activity and standard counting errors were recorded for each measurement. For the paired petri dish measurements, the ratio and mean were calculated. Estimated aliquot activity concentrations were based on the mean. In the same way, the overall sample standard deviation and % CV were calculated. For locations with samples collected at multiple depths, ratios of the respective mean sample activity concentrations were calculated.
- **6. University of Pittsburgh Work.** Capt Craig Refosco, an Air Force active duty health physicist, performed additional analysis of select samples in partial fulfillment of the requirements for the M.S. in Science Degree (Refosco 2001). Five samples were used for the analysis. Smaller aliquots were prepared by subdivision of 200 g aliquots into ten 20 g aliquots. The samples were placed in the lids of same type of petri dish that was used by AFIERA, enclosed by the petri dish bottom portion, and sealed with silicone. The aliquots were counted on a Bicron (12.7 cm diameter x 1.6 mm thick) NaI(Tl) and calibrated with an AFIERA prepared NIST traceable source.

Results

1. Sample Activity Concentrations. Table A in Appendix A contains the analytical results from the laboratory γ -spectroscopy. The reported $^{239/240}$ Pu activity concentration for the individual aliquots is based on assessment of the 241 Am 60 keV γ -ray and an assumed $^{239/240}$ Pu to 241 Am ratio of 5.4 (Rademacher 1999a). Uncertainty values are based on randomness due to counting statistics and are expressed at the 95 % confidence interval (CI). Aliquot $^{239/240}$ Pu activity concentrations, estimated from individual aliquot measurements (one side alone), ranged from 1.1 to 68,500 pCi g⁻¹. For the mean of paired aliquot measurements, $^{239/240}$ Pu activity concentrations ranged from 1.1 to 51,700 pCi g⁻¹.

2. Heterogeneity.

- a. General. This study provides a description of heterogeneity at four levels. First, for seven sampling locations, surface soil samples within 30 cm of the central location describe variability within sampling regions. Second, the evaluation of heterogeneity at the sample level is described through multiple aliquot analysis of samples. Third, conjugate measurement of individual aliquots provides examination of heterogeneity at the aliquot level. Lastly, work performed at the University of Pittsburgh, describes heterogeneity at the sub-aliquot level of 20 g.
- b. Heterogeneity at Sampling Level. Table 4 contains ^{239/240}Pu activity concentrations for the surface soils of sample locations 1 through 7. For each location, results are provided for the center, west, east, north, and south sub-sample locations. The last column lists the variability of the measurements expressed in % CV. The variability among the locations ranges from 32 to 186 %. Sampling location 1 had the greatest level of variability, with the most extreme range of activity concentration being 3.2 to 2,792 pCi g⁻¹. The other three samples collected at this location had activity concentration clustered around 140 pCi g⁻¹. Sampling location 7 had the highest mean activity concentration among the sub-samples, being 9,647 pCi g⁻¹. For this sampling location, the ratio between the highest and the lowest activity concentration among sub-samples was about 4, while that of sample location 1 was almost 1,000.

Table 4. Sub-Sample ^{239/240}Pu Activity Concentrations.

Sample		Variability					
Location	Center	W	Е	N	S	Mean	(% CV)
1	2,792.2	165.4	127.3	3.2	138.6	645	186
2	594.1	268.8	184.2	354.9	553.5	391	46
3	232.8	180.9	251.1	98.6	173.5	187	32
4	1,004.8	1,227.3	437.9	774.6	1,049.6	899	34
5	577.5	278.8	572.3	543.6	199.8	434	42
6	2,207.1	75.9	885.3	2,560.7	4,681.5	2,082	85
7	5,273.2	12,888.2	19,889.6	3,866.5	6,319.3	9,647	69

c. Heterogeneity at Sample and Aliquot Level.

Figure 4 contains a histogram of the individual aliquot data from Table A-1 for sample 14 (abbreviated notation based on the last digits of the base sample number). Apparent is the vast discrepancy among individual aliquot activity concentrations. The mean aliquot activity concentrations are: 544, 7,742, and 90 pCi g⁻¹, respectively for increasing aliquot number. The ratio between mean activity concentration of aliquots 2 and 3 is 86. With the extensive blending and homogenizing procedures applied to the samples during preparation, heterogeneity of this magnitude was not anticipated. While the composition of an individual aliquot cannot be inferred from the data presented here, it is clear that in the case of aliquot 2, that the majority of sample activity is concentrated in a single or possibly a few particles. If the activity is primarily concentrated in a single particle, the minimum particle diameter is $162 \mu m$, under the assumption that the plutonium is in a dioxide form ($\rho = 9.8 \text{ g cm}^{-3}$). Regardless of the particle(s) size(s) and composition, the ratio of the estimated activity concentration from the down to up measurements is 4.3, and this discrepancy may introduce significant errors if the sample was measured in one configuration only.

Figure 4. Sample 14 $^{239/240}$ Pu Activity Concentration for Individual Aliquots (Mean = 2792 pCi g⁻¹, % CV = 176 %).

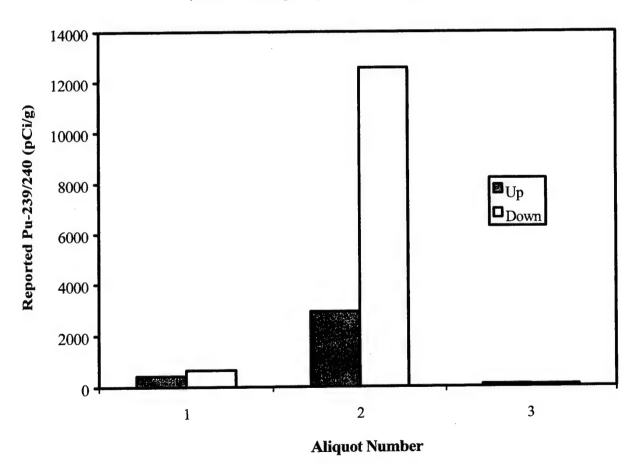


Figure B-1 of Appendix B contains a histogram of the individual aliquot data from Table A for sample 67. Apparent from the histogram, aliquots 1, 3, and 5 have relatively consistent activity concentration, while 2 and 4 are considerably higher. The mean aliquot activity concentrations are: 92, 607, 101, 500, and 55 pCi g⁻¹, respectively for increasing aliquot number. For aliquots 2 and 4, the ratio of down to up is 1.7 and 1.4, respectively. Because of the high sample activity concentration of these aliquots, the high conjugate counting ratios are indicative of heterogeneity effects rather than uncertainties of counting statistics. For these aliquots, if the activity is largely associated by single particles, the minimum particle diameters would be 69 and 65 μm, respectively (dioxide form of plutonium). For aliquots 1, 3, and 5, the plutonium activity appears to be relatively homogenous in distribution, based on agreement between respective conjugate measurements.

Figure B-2 contains a histogram of the individual aliquot data from Table A for sample 15. Like sample 14, one aliquot (2) contains significantly higher activity concentration than the others. The disparity between the up and down measured activity concentration, 4.6, is indicative of a single particle dominating the aliquot activity. Figure B-3 contains a histogram of the individual aliquot data for sample 53. Like samples 14 and 15, this sample has one aliquot that contains a significantly higher mean activity concentration than the others. Sample 53 is unique to this data set in having one of the highest individual aliquot mean activity concentrations, 33,200 pCi g⁻¹. Like many of the other high activity concentration aliquots, the ratio of the conjugate measurements is high with a value of 2.6. For this aliquot, if the activity concentration is concentrated in an individual particle, the minimum particle diameter is 260 μm (dioxide form of plutonium).

The samples discussed above provide a unique description of heterogeneity of WGP and represent the greatest degree of heterogeneity among those evaluated in this study. Many of the samples had less apparent heterogeneity. Figure B-4 contains a histogram of the individual aliquot data for sample 75. From the plot, it is apparent that close agreement exists among the conjugate aliquot measurements and reasonably well among all measurements. The % CV for the individual measurements was 72 %. Figure B-5 contains a histogram of the individual aliquot data for sample 60. This sample has a mean activity concentration of 128 pC g⁻¹ and a % CV of only 13 %. The variability among aliquots and conjugate measurements is low, suggesting a homogenous sample.

Figure 5 contains a scatterplot of % CV of individual aliquot measured ^{239/240}Pu activity concentration versus the mean sample activity concentration. The plot describes the heterogeneity that exists among individual aliquots of a sample. Samples with the highest variability among the measurements are those with the highest mean activity concentration. For samples with mean activity concentration between 100 and 1,000 pCi g⁻¹, the % CV values range from very low values to about 160 %. Thirteen samples have mean activity concentrations between 10 - 100 pCi g⁻¹, and have % CV values ranging from 18 to 128 %. Seven samples have mean activity concentrations less than 10 pCi g⁻¹, and have % CV values ranging from 25 to 137 %. The samples in this range of activity concentration have particular significance to the BOMARC site because the remediation criterion for the site is 8 pCi g⁻¹.

Figure 6 contains a scatterplot of the ratio of the conjugate measurements of ^{239/240}Pu activity concentrations for individual aliquots (reciprocal plotted for ratios less than 1). The plot describes the heterogeneity that exists within individual aliquots. In general, samples with the highest ratios are those with the highest mean ^{239/240}Pu activity concentration. The ratios ranged from 1.0 to 4.6,

Figure 5. Percent Coefficient of Variation (% CV) of Individual Aliquot Measured ^{239/240}Pu vs. Mean Sample ^{239/240}Pu.

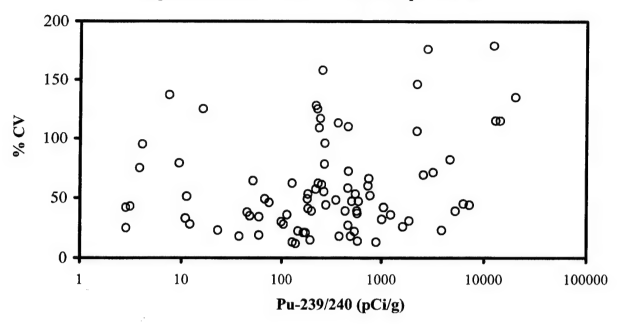
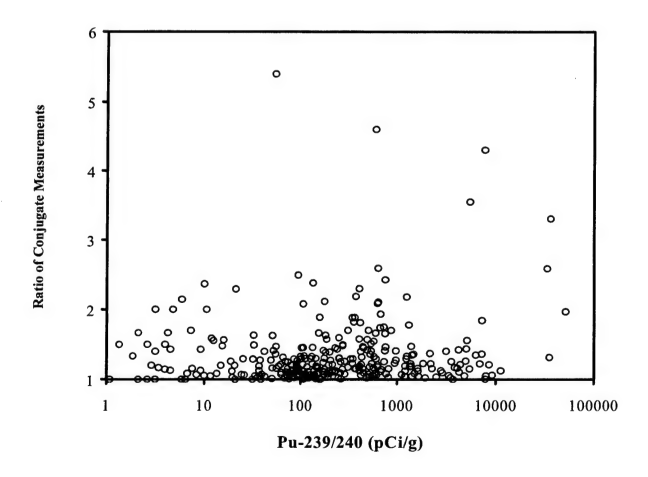


Figure 6. Ratio of Conjugate Measurements of ^{239/240}Pu for Individual Aliquots.



with the majority being below 3. In general, higher ratios are indicative of a greater degree of heterogeneity, indicating that the highest heterogeneity exists among samples of the highest activity concentration. Many of the samples have ratios near one. It could be argued in general that these samples have a low degree of heterogeneity. On an individual sample basis, however, conclusions cannot be made, due to the potential for single or multiple particle samples having the particle(s) on the mid-line of the sample. The effect of heterogeneity was exhibited for samples across entire range of activity concentration.

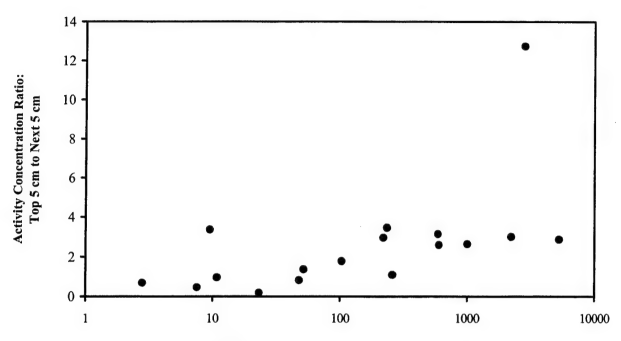
3. University of Pittsburgh Sub-Aliquot Data.

Samples: 50, 51, 52, 57, and 79 had 10 sub-aliquots analyzed from the first aliquot. The results of the analysis are listed in Table B-3 with the results from the AFIERA analysis. For each set of sub-aliquots, the mean, % CV, and relative percent difference (RPD) between the mean and AFIERA result. For four of the sub-aliquot groups, the degree of heterogeneity observed among the sub-aliquots was low, with the maximum to minimum sub-aliquot ratios ranging from 2.9 to 4.3 and range of % CV values ranging from 32 to 50. For these samples, it was initially assumed that the contaminant was uniform with the ratio of the AFIERA conjugate measurements having ratios less than 1.17. Further, for samples 50, 51, and 52, the entire samples overall had a low degree of heterogeneity, with combined aliquot % CV values of 39, 31, and 14. For sample 57, the combined aliquot % CV was 71, but the greatest degree of heterogeneity within this sample was expected in aliquot 2, with a conjugate measurement ratio of 1.36, as compared to 1.15 in aliquot 1 (the aliquot that was sub-divided).

The aliquot from sample 79 exhibited the greatest degree of heterogeneity among the five samples with a % CV among the sub-aliquots at 306 %. The first sub-aliquot contained over 97 % of the total aliquot activity and had an activity concentration 800 times the lowest activity sub-aliquot. If all the activity for this sub-aliquot is comprised of a single particle, the minimum diameter is 250 µm. Another interesting observation with this aliquot is the existence of a sub-aliquot with an intermediate activity concentration: 12.7 times that of the lowest, but 63 times lower than the highest. For the remaining sub-aliquots, the activity concentration was fairly uniform.

4. Depth Distribution. Seventeen locations had samples collected at the 5-10 cm depth. For one location, another sample was collected at a depth of 10-15 cm. The ratio of the $^{239/240}$ Pu activity concentration in the top 5 cm to the next 5 cm is plotted in Figure 7. The ratios range from 0.2 to 12.7. While the ratio among paired samples has a significant degree of variability, among the higher activity concentration samples, better agreement existed, with the median being about 3. For all paired samples, the median ratio is 2.6. The sample with the highest ratio is comprised of samples 14 and 15 (14 being the top 5 cm). Sample 14 (see Figure 4) has a % CV of 176 % among its aliquots. This value is second highest among the sample evaluated. As well, this sample was in the group of samples that exhibited a high degree of spatial heterogeneity (see Table 4, sampling location #1). The unusually high heterogeneity exhibited in this sample may in part explain non-agreement with the depth distribution of the other high activity concentration samples.

Figure 7. Activity Concentration Ratio: Top 5 cm to Next 5 cm vs. ^{239/240}Pu Activity Concentration in Top 5 cm.



Pu-239/240 Activty Concentration in Top 5 cm (pCi/g)

5. FIDLER Correlation Coefficients.

The correlation between concentration of WGP in soils and the response of portable γ -radiation instruments is important in assessing instrument minimum detectable concentrations (MDCs). The FIDLER instrument is the most common for in-situ γ - radiation measurements of WGP contaminated soils. Figure 8 contains a scatterplot of FIDLER response versus the mean ^{239/240}Pu activity concentration in the top 5 cm of soil of those from the first sampling effort. Apparent from the plot is a high degree of variability in the data set, based on the low squared correlation coefficient (R²) of 0.35. The intercept of the linear regression was set at 1,000 cpm, the background count rate of the FIDLER in an area of the site where WGP is known not to have been deposited.

The regression analysis is affected to a significant degree by the two highest activity concentration data points. Another regression of the data is provided in Figure 9, with the two highest activity concentration data points truncated from the plot. Visually, the data are in better agreement with the regression line and is reflected in a significantly higher R² of 0.84. The slope of the regression analysis was increased by a factor of 2.3 over that of the complete data set. Table 5 provides a list of estimated correlation coefficients of FIDLER response and ^{239/240}Pu activity concentration for the BOMARC site, based on an assumed activity concentration ratio between 5 cm successive lifts of 2.6.

Figure 8. Scatterplot of FIDLER Response vs. $^{239/240}$ Pu Activity Concentration (Full Data Set, n = 30)

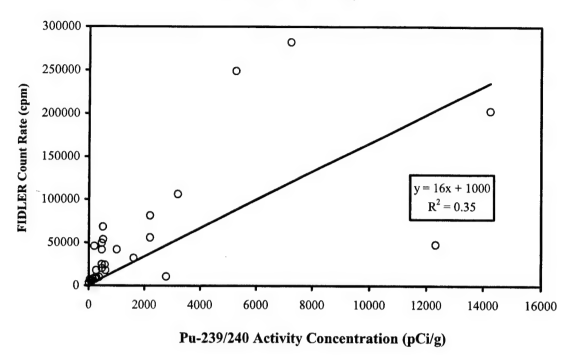


Figure 9. Scatterplot of FIDLER Response vs. $^{239/240}$ Pu Activity Concentration (Truncated Data Set, n = 28)

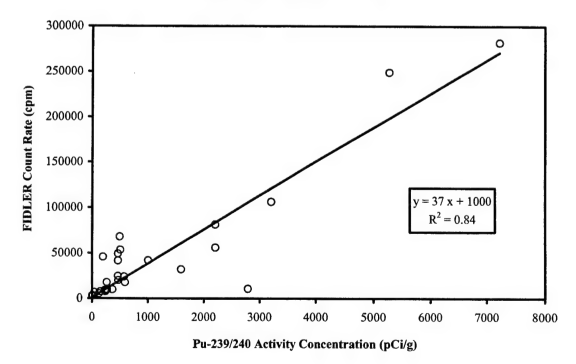


Table 5. Estimated Correlation Coefficients of FIDLER Response and ^{239/240}Pu Activity Concentrations.

Averaging Depth*	Correlation Coefficient
(cm)	(cpm g pCi ⁻¹)
5	37
10	53
15	72

^{*} Assummed ratio of ^{239/240}Pu between successive 5 cm lifts is 2.6.

Discussion and Conclusions

Heterogeneity impacts laboratory evaluation of soil samples when γ -radiation evaluation of the 60 keV photon from ²⁴¹Am is used with an assumed constant ratio between ²⁴¹Am and ^{239/240}Pu. Heterogeneity was observed at every level of evaluation: sub-aliquot, aliquot, sample, and sampling location. The conjugate counting methodology used aided in assessing the degree of heterogeneity. For many aliquots evaluated, the conjugate counting methodology improved measurement accuracy. For WGP samples, the conjugate counting methodology should be implemented unless homogeneity can be verified.

Due to the heterogeneity among aliquots from the same sample, the conjugate method should be modified to accommodate larger samples. This requirement necessitates larger diameter detectors. HpGe detectors are not commonly manufactured with diameters in excess of 10 cm. NaI(Tl) detectors are a reasonable alternative. AFIERA has purchased a pair of 20 cm diameter x 2 mm thick NaI(Tl) detectors. Over 2,000 g of soil can be held in sample containers of the same diameter and 4.5 cm in height.

Heterogeneity impacts the accuracy of the estimated correlation coefficient between the FIDLER response and $^{239/240}$ Pu activity concentrations. Clear from the evaluation provided here, careful review of the data and accurate soil measurement techniques should be used to estimate this parameter. Evaluation of vertical distribution of the contaminant is critical to in-situ γ -measurements if they are being used to meet remediation or screening criteria. For the evaluation provided here, there was good agreement in 10 of the 17 soil samples evaluated for vertical distribution of the contaminant.

Acknowledgements

Mr. Dale Thomas - Laboratory Analysis Oversight and Data Review

Major Kevin Martilla - 1999 Soil Sampling Effort

Capt Eugene Sheely – 2000 Soil Sampling Effort

TSgt Ty Richards, Laboratory Sample Analysis

Mr. James Sablan - Laboratory Sample Preparation

Mr. William Walker - Laboratory Sample Analysis

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Appendix A Laboratory ^{239/240}Pu Analysis Results

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Table A-1. Individual Aliquot 239/240Pu Activity Concentrations. (1999 Sampling)

		1		Г	T	Τ	1	ι			1	_	_		_		1		r				1		г —		
	Ci/g)	HIGH	LOW	4.31	4.60	1.12	2.11	1.33	1.07	1.75	2.42	1.70	1.01	1.70	1.88	1.12	2.08	1.30	1.82	1.33	1.11	1.46	1.23	1.09	1.09	1.21	1.78
	ntration (r	vn	Uncer.	829	232	42	56	0.5	30	217	103	150	38	92	106	51	34	99	109	35	38	145	06	336	1467	162	221
Aliquot #2	vity Conce	Down	Value	2920	1,000	179	242	1.6	131	935	444	648	165	395	456	221	144	286	467	150	189	623	387	1,450	633	869	952
<i>H</i>	Pu-239/240 Activity Concentration (pCi/g)	d	Uncer.	2,920	50	37	27	0.5	59	124	250	68	38	54	99	58	16	51	59	27	39	212	73	365	135	135	393
	Pu-23	Up	Value	12,570	217	160	115	2.2	123	535	1,080	381	164	233	243	248	69	220	257	113	170	911	315	1,570	581	578	1,690
	Ci/g)	HIGH	LOW	1.51	1.11	1.57	1.10	1.40	1.17	1.41	1.45	1.15	1.59	1.36	1.65	1.05	1.36	1.12	1.07	1.19	1.39	1.29	1.32	1.36	1.47	1.33	1.08
	40 Activity Concentration (pCi/g)	wn	Uncer.	101	47	54	18	0.5	38	205	20	39	45	105	133	22	15	24	34	19	55	325	88	395	845	251	187
Aliquot #1	vity Conce	Down	Value	434	203	230	80	2.7	162	884	87	167	196	453	571	95	63	103	145	83	235	1,400	379	1,700	366	1,080	208
		Up	Uncer.	152	52	34	17	1.1	32	146	29	45	72	143	219	21	11	27	32	23	39	252	116	290	125	333	174
	Pu-239/2	_	Value	655	226	146	73	3.8	138	679	126	192	312	617	945	06	46	115	136	99	169	1,080	200	1,250	536	1,430	748
ling	tion	Sub-	sample	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S
Sampling	Location	#	1			-	-	•		2					w						4						
0	Base	Number	GS0000014 GS0000015 GS0000016 GS0000018 GS0000019 GS0000020 GS0000021 GS0000024 GS0000025 GS0000026 GS0000026 GS0000027 GS0000027 GS0000027 GS0000028 GS0000028 GS0000033 GS0000033								GS0000035	GS0000036	GS0000037														

Table A-1. Individual Aliquot ^{239/240}Pu Activity Concentrations (Continued) (1999 Sampling)

Aliquot #4	Pu-239/240 Activity Concentration (pCi/g)	Down HIGH	Value Uncer. LOW				65 15 1.25	15	15	15	15	15	15	15	23	15	23	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6	23 23 8.6
Up	Up	T 14000	Oncer.		19						20 1						8.1										
	Pu-2		Value		82						98						36	:									
	oCi/g)	HIGH	TOW	1.13	1.22	1.05		1.67		1.26	1.11	1.04	2.38	1.27		1.88	1.16			1.15	1.12		1.29	1.14	1.30	1.40	
	entration (Down	Uncer.	22	23	32		0.5		75	22	26	1.1	42		09	1			18	35		93	173	89	116	
Aliquot #3	240 Activity Concentration (pCi/g)	Do	Value	96	100	136		3.2		324	92	1111	80	182		258	47			78	147		399	744	289	501	
	9/240 Acti	þ	Uncer.	19	19	33		1.1		09	24	27	44	54		113	13			16	31		72	152	52	83	
	Pu-239/	$\Omega_{\mathbf{p}}$	Value	84	82	143		5.4		258	102	115	190	232		485	55			89	131		309	653	222	359	
Sampling	ation	-qnS	sample	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S
Sam	Sampling Location Sul					-	-			2							7	C			4						
Desc	Sample	Number	Iduilibei	GS0000014 GS0000015 GS0000016 GS0000017 GS0000018					GS0000020	GS0000021	GS0000022	GS0000023	GS0000024	GS0000025	GS0000026	GS0000027	GS0000028	GS0000029	GS0000030	GS0000031	GS0000032	GS0000033	GS0000034	GS0000035	GS0000036	GS0000037	

Table A-1. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (1999 Sampling)

Sam	Sampling			Aliquot #5					Aliquot #6		,
Loc	Location	Pu-239/2	9/240 Acti	vity Conc	40 Activity Concentration (pCi/g)	oCi/g)	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	vity Conc	entration (J	Ci/g)
#	-qnS	ח	Up	Do	Down	HIGH	Ω	Up	oQ	Down	HIGH
t !	sample	Value	Uncer.	Value	Uncer.	TOW	Value	Uncer.	Value	Uncer.	LOW
	0-2 C										de la la companya de
	2-4 C	106	25	112	56	1.06					
-	0-2 W										
-	0-2 E										
	0-2 N										
	0-2 S										
	0-2 C										
	2-4 C	140	32	163	38	1.16	207	48	110	25	1.89
c	0-2 W										
7	0-2 E										
	0-2 N										
	0-2 S										
	0-2 C										
	2-4 C	84	19	62	23	1.15					
C	0-2 W										
C	0-2 E										
	0-2 N										
	0-2 S										
	0-2 C										
	2-4 C										
_	0-2 W										
†	0-2 E										
	0-2 N										
	0-2 S										0

Table A-1. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (1999 Sampling)

		ficient	ion																								
Aliquot	arameters	Percent Coefficient	Of Variation	176	128	21	62	43	12	47	125	78	41	48	53	62	49	49	61	30	21	32	18	36	39	52	42
Combined Aliquot	Statistical Parameters	Mean Activity	Concentration (pCi/g)	2,790	219	165	127	3.2	139	594	228	269	184	352	554	233	89	181	251	66	174	1,000	381	1,230	438	775	1,050
ling	tion	-qnS	sample	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S
Sampling	Location	#	-			-	-					C	7					,	C					_	†		
Base	Commit	Number		GS0000014	GS0000015	GS0000016	GS0000017	GS0000018	GS0000019	GS0000020	GS0000021	GS0000022	GS0000023	GS0000024	GS0000025	GS0000026	GS0000027	GS0000028	GS0000029	GS0000030	GS0000031	GS0000032	GS0000033	GS0000034	GS0000035	GS0000036	GS0000037

Table A-1. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (1999 Sampling)

Sample		J Sumdina		7	wildnor #1				7	7# lonbirs		
	Loca	Location	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	vity Conce	entration (1	Ci/g)	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	ivity Conce	entration (1	Ci/g)
	#	-qnS	n	Up	Do	Oown	H9IH	dU	۵	Po	Down	HIGH
Number	ŧ	sample	Value	Uncer.	Value	Uncer.	LOW	Value	Uncer.	Value	Uncer.	LOW
GS0000038		0-2 C	629	153	825	192	1.25	321	75	505	117	1.58
GS0000039		2-4 C	234	55	293	89	1.25	84	19	122	28	1.45
GS0000040	V	0-2 W	274	64	251	58	1.09	239	99	523	122	2.19
GS0000041	٠	0-2 E	609	142	388	90	1.57	416	26	875	203	2.10
GS0000042		0-2 N	481	112	417	97	1.15	989	136	069	160	1.18
GS0000043		0-2 S	268	62	566	62	1.01	135	31	130	30	1.04
GS0000044		0-2 C	2,420	562	8,590	1,990	3.55	492	114	5023	117	1.02
GS0000045		2-4 C	393	91	476	111	1.21	1,630	379	1,210	281	1.35
GS0000046	9	0-2 W	1,720	399	789	183	2.18	393	16	441	103	1.12
GS0000047	>	0-2 E	849	197	1,040	243	1.23	881	205	992	178	1.15
GS0000048		0-2 N	1,260	293	1,500	349	1.19	2,480	929	2,570	965	1.03
GS0000049		0-2 S	1,240	288	1,520	353	1.23	8,180	1,900	1,790	1,810	1.05
GS0000050		0-2 C	4,800	1,110	3,960	920	1.21	3,980	923	3,170	136	1.25
GS0000051		2-4 C	2,640	612	2,960	687	1.12	1,970	457	1,930	447	1.02
GS0000052	1	4-6 C	534	124	623	145	1.17	849	149	720	167	1.11
GS0000053		0-2 W	5,370	1,250	7,240	1,680	1.35	6,240	1,450	7,630	1,770	1.22
GS0000054		0-2 E	4,500	1,040	4,080	946	1.10	34,800	8,090	68,470	15,900	1.96
GS0000055		0-2 N	4,160	996	3,560	826	1.17	2,760	642	3,020	702	1.09
GS0000056		0-2 S	7,780	1,810	9,400	2,180	1.21	3,280	160	3,460	803	1.06
GS0000057	∞	0-2	2,440	267	2,120	493	1.15	8,360	1,940	6,150	1,430	1.36
GS0000058	6	0-2	9.8	2.2	2.6	2.2	1.13	11	2.7	12	2.7	1.05
GS0000059	10	0-2	3.8	1.1	2.7	0.5	1.40	2.2	0.5	1.6	0.5	1.33
GS0000060	11	0-2	133	31	157	37	1.18	126	56	114	27	1.11
GS0000061	12	0-2	2.2	0.5	1.6	0.5	1.33	3.8	1.1	8.1	1.6	2.14

Table A-1. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (1999 Sampling)

Aliquot #3 Pu-239/240 Activity Concen Up Dow	HS W	Pu-239/240 Ac Up Value Uncer.	Aliquot #4 Pu-239/240 Activity Concentration (pCi/g)	(pCi/g) HIGH LOW
0-2 C 2-4 C 0-2 W 200 46 18	186 43 1.07			
0-2 S 604 140 65	639 148 1.06			
551 128	135 1.05	535 124	515 110	1.04
0-2 W 504 117 60	666 155 1.32			
0-2 N 6,240 1,453 4,0	4,010 932 1.56 1,3	1,310 305	1,100 256	1.19
0-2 C 7,880 1,830 7,850	50 1,820 1.00			
1,230 285	274 1.04	2,510 583	1,830 426	1.37
496 115	124 1.07		536 124	1.29
0-2 W 47,900 11,100 18	18,600 4,320 2.58 5,9	5,970 1,390	4,160 967	1.43
4,730 1,200	537			
2,200	2,080 1.06	4,400 1,020	3,790 881	1.16
0-2 3,680 854 2,6	2,630 610 1.40 1,5	1,550 359	1,730 402	1.12
0-2 12.4 3.2 18	18.4 4.3 1.48			
0-2 5.4 1.1 3.	3.2 0.5 1.67			
0-2 120 25 1	146 34 1.22 10	109 25	117 27	1.08
0-2	34 1.22			

Table A-1. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (1999 Sampling)

Aliquot #6	Pu-239/240 Activity Concentration (pCi/g)	Up Down HIGH	Uncer. Value Uncer.														90 323 1,450 337 1.04	5 106 639 149 1.40									
		HIGH	LOW Value														1.02 1,390	1.05 455					1.06				
	40 Activity Concentration (pCi/g)		Uncer.														371	127					382				
Aliquot #5	ivity Conce	Down	Value														1,600	549					1,650				
	39/240 Act	Up	Uncer.		-												363	134					406				
	Pu-239/2	7	Value														1,570	278					1,750				
Sampling	Location	-qnS	sample	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	4-6 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2	0-2	0-2	0-2	
Sam	Loc	7	‡			ų	<u>0</u>					7	5					7	`				8	6	10	11	
	Base	Sample	Number	GS0000038	GS0000039	GS0000040	GS0000041	GS0000042	GS0000043	GS0000044	GS0000045	GS0000046	GS0000047	GS0000048	GS0000049	GS0000050	GS0000051	GS0000052	GS0000053	GS0000054	GS0000055	GS0000056	GS0000057	GS00000058	GS0000059	GS0000060	

Table A-1. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (1999 Sampling)

Aliquot arameters	Percent Coefficient	37	53	44	39	22	39	146	09	99	13	69	82	39	31	14	115	135	23	45	71	28	43	13	75
Combined Aliquot Statistical Parameters	Mean Activity Concentration (nCi/a)	578	183	279	572	544	200	2,220	737	752	885	2,560	4,680	5,270	1,850	584	12,900	19,900	3,870	6,320	3,210	12	3.2	128	3.9
oling tion	Sub- sample	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2 C	2-4 C	4-6 C	0-2 W	0-2 E	0-2 N	0-2 S	0-2	0-2	0-2	0-2	0-2
Sampling Location	#		•	v	ე					7)						7				8	6	10	11	12
Base	Sample Number	GS0000038	GS0000039	GS0000040	GS0000041	GS0000042	GS0000043	GS0000044	GS0000045	GS0000046	GS0000047	GS0000048	GS0000049	GS0000050	GS0000051	GS0000052	GS0000053	GS0000054	GS0000055	GS0000056	GS0000057	GS0000058	GS0000059	GS0000060	GS0000061

Table A-1. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (1999 Sampling)

		_		_				,	_				,	_	,		,				
	Ci/g)	HIGH	LOW	1.26	1.63	1.02	1.04	1.18	1.66	1.18	1.42	1.26	1.70	1.04	1.15	2.09	1.28	1.06	1.81	1.09	1.15
	ntration (r	wn	Uncer.	24	15	25	394	247	106	132	1,150	1,290	254	45	83	96	253	103	127	45	81
Aliquot #2	vity Conce	Down	Value	103	63	109	1,700	1,520	456	572	4,960	5,550	1,100	191	355	412	1,090	443	545	196	346
7	Pu-239/240 Activity Concentration (pCi/g)	d	Uncer.	19	9.2	25	378	139	176	113	810	1,020	150	46	95	200	199	26	70	50	93
	· Pu-23	$^{ m Up}$	Value	82	39	107	1,630	1,280	757	486	3,490	4,400	645	198	408	098	854	419	301	213	399
	Ci/g)	HIGH	LOW	1.74	1.19	1.38	1.07	1.16	1.13	1.54	1.31	1.15	1.01	1.14	1.03	1.07	1.13	1.23	1.07	1.21	3.30
	Pu-239/240 Activity Concentration (pCi/g)	Down	Uncer.	115	12	46	251	351	20	106	7,030	1,350	24	69	188	65	54	116	42	55	12,900
Aliquot #1	ivity Conc	Do	Value	493	54	212	1,080	1,510	86	457	30,300	5,830	105	294	809	281	231	498	181	237	55,500
	9/240 Act	Up	Uncer.	199	15	36	268	406	23	164	9,230	1,180	25	60	193	70	48	142	39	45	3,900
	Pu-23	ח	Value	856	64	153	1,150	1,750	97	706	39,800	5,070	106	257	832	302	205	611	170	196	16,800
Sampling	ation	Sub-	sample	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sam	Location	#	E	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30
Dege	Dase	Number	INUITION	GS0000062	GS0000063	GS0000064	GS0000065	GS0000066	CS0000067	GS0000068	GS0000069	GS0000070	GS0000071	GS0000072	GS0000073	GS0000074	GS0000075	GS0000076	GS0000077	GS0000078	GS0000079

Table A-1. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (1999 Sampling)

											Γ										
	oCi/g)	HIGH	TOW	1.29		1.03	1.23	1.22	1.42	1.19				1.20	1.10		1.45	1.04			
	entration (1	Down	Uncer.	27		32.9	473	461	96	68				42	56		44	139			
Aliquot #4	vity Conce	Do	Value	115		142	2,040	1990	414	382				179	241		189	599			
	Pu-239/240 Activity Concentration (pCi/g)	d	Uncer.	21		34	385	564	136	105				50	51		64	134			
	Pu-23	Up	Value	68		146	1,660	2,430	587	454				216	218		274	577			
	Ci/g)	HIGH	TOW	1.21	1.06	1.08	1.41	1.04	1.33	1.29	1.02	1.12	1.29	1.02	1.08		1.45	1.17	1.12	1.00	1.04
	intration (p	wn	Uncer.	22	6.5	33	264	279	27	88	807	2,460	34	57	133		126	103	43	37	113
Aliquot #3	vity Conce	Down	Value	96	27	143	1,140	1,200	115	377	3,470	10,600	1,450	247	572		542	443	187	160	486
7	Pu-239/240 Activity Concentration (pCi/g)	Up	Uncer.	27	5.9	36	188	586	20	89	826	2,760	26	59	143		87	88	49	37	117
	Pu-23	n	Value	116	25	155	608	1,250	98	292	3,560	11,900	112	253	617		373	380	500	160	503
ling	tion	-qnS	sample	0-5	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sampling	Location	#	ŧ	13	14	15	16	17	18	16	20	21	22	23	24	25	26	27	28	29	30
0.00	base	Number	Number	GS0000062	GS0000063	GS0000064	GS0000065	GS0000066	GS0000067	GS0000068	6900000SD	0200000SD	GS0000071	GS0000072	GS0000073	GS0000074	GS0000075	GS0000076	GS0000077	GS0000078	GS0000079

Table A-1. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (1999 Sampling)

San	Sampling			Aliquot #5					Aliquot #6		
현	Location	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	ivity Conce	entration (1	oCi/g)	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	ivity Conc	entration (1	Ci/g)
	-qnS	ח	Up	Do	Down	HIGH	7	Up	Do	Down	НІСН
	sample	Value	Uncer.	Value	Uncer.	row	Value	Uncer.	Value	Uncer.	LOW
	0-2										
,	0-2										
	0-2										
16	0-2	9,430	2,190	5,130	1,190	1.84	1,240	287	1,460	339	1.18
17	0-2										
18	0-2	45	10	99	15	1.47					
19	0-2										
20	0-2										
21	0-2										
22	0-2										
23	0-2	1,660	2988	1,130	262	1.47					
24	0-2										
25	0-2										
26	0-2										
27	0-2										
28	0-2										
29	0-2										
30	0-2										

Table A-1. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (1999 Sampling)

_				_	_	1					1			_	_						_
Aliquot	arameters	Percent Coefficient	Of Variation	117	38	22.3	106	26	96	27	115	44	113	110	47	58	72	18	55	15	179
Combined Aliquot	Statistical Parameters	Mean Activity	Concentration (pCi/g)	244	46	146	2,210	1,610	271	466	14,300	7,230	368	463	507	464	470	496	266	194	12,300
	Ci/g)	HIGH	TOW				1.12														
	entration (r	wn	Uncer.				566														
Aliquot #7	Pu-239/240 Activity Concentration (pCi/g)	Down	Value				1,140														
7	9/240 Acti	Ď	Uncer.				297														
	Pu-23	Up	Value				1,280														
Sampling	ıtion	Sub-	sample	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	2-0
Sam	Location	#	#	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30
þ	Base	Sample	Ivalliber	GS0000062	GS0000063	GS0000064	GS0000065	GS0000066	GS0000067	GS0000068	6900000SD	GS0000070	GS0000071	GS0000072	GS0000073	GS0000074	GS0000075	GS0000076	GS0000077	GS0000078	GS0000079

Blank Cells Represent Samples Where an Aliquot Did Not Exist (Aliquot Number Varied from 2 to 10 per Sample)

Table A-2. Individual Aliquot ^{239/240}Pu Activity Concentrations. (2000 Sampling)

	Sam	Sampling			Aliquot #1					Aliquot #2		
Base	Loc	Location	Pu-239/2	9/240 Acti	vity Conce	40 Activity Concentration (pCi/g)	Ci/g)	Pu-23	Pu-239/240 Activity Concentration (pCi/g)	vity Conce	entration (p	Ci/g)
Sample	#	Sub-	U	Up	Do	Down	HIGH	Up	p	Do	Down	HIGH
Number	‡	sample	Value	Uncer.	Value	Uncer.	LOW	Value	Uncer.	Value	Uncer.	LOW
6990000SD	31	2-4	49	11	35	8.1	1.40	39	9.2	39	9.2	1.00
GS0000670	32	2-4	154	36	161	44	1.24	93	22	125	29	1.34
GS0000671	33	2-4	61	14	65	15	1.07	75	17	96	22	1.29
GS0000672	34	2-4	41	6.7	38	9.8	1.07	117	27	111	26	1.05
GS0000673	35	2-4	3.8	1.1	2.7	0.5	1.40	2.7	0.5	2.7	6.0	1.00
GS0000674	36	2-4	14	3.2	16	3.8	1.20	21	4.9	21	4.9	1.00
GS0000675	37	2-4	50	12	51	12	1.01	2.7	0.5	3.8	1.1	1.40
GS0000676	38	2-4	2.7	0.5	1.6	0.5	1.67	1.1	0.4	1.1	0.5	1.00
GS0000677	39	2-4	99	15	09	14	1.10	57	13	75	11	1.31
GS0000678	40	2-4	465	108	668	208	1.93	150	35	161	37	1.08
6290000SD	31	0-2	55	13	43	10	1.28	135	31	54	12	2.49
GS0000680	32	0-2	32	9.7	32	9.7	1.00	21	4.9	18	4.3	1.12
GS0000681	33	0-2	69	16	89	16	1.02	35	39	39	9.2	1.12
GS0000682	34	0-2	200	46	286	99	1.43	108	25	109	25	1.01
GS0000683	35	0-2	5.4	1.1	9.2	2.2	1.70	3.8	1.1	5.4	1.1	1.43
GS0000684	36	0-2	6.7	2.2	10	2.7	1.06	13	3.2	14	3.2	1.08
GS0000685	37	0-2	2.2	0.5	2.2	0.5	1.00	3.8	1.1	4.3	1.1	1.14
GS0000686	38	0-2	3.2	1.1	3.2	1.1	1.00	1.6	0.5	2.2	0.5	1.33
GS0000687	39	0-2	109	25	159	37	1.46	77	18	82	19	1.07
GS0000688	40	0-2	223	52	333	77	1.50	145	34	144	34	1.01

Table A-2. Individual Aliquot 239/240Pu Activity Concentrations (Continued) (2000 Sampling)

						-	_	-	-y							, .							
	Ci/g)	HIGH	MOI		1 07	1 19		1.50	100	2.29	2.00	1.16	1.66	1.21	1.07	1.04	1.09	1.59	1.43	1.08	1.17	1.18	1.41
	intration (r	wn	Uncer.		22	9.2	!	0.5	1.6	7.0	1.6	14	45	16	5.9	8.6	112	3.2	1.6	1.6	0.5	25	47
Aliquot #4	vity Conce	Down	Value		96	41		1.6	6.5	30	6.5	09	194	89	25	37	481	15	7.6	6.5	3.2	108	203
7	Pu-239/240 Activity Concentration (pCi/g)	d	Uncer.		21	8.1		0.4	1.6	3.2	1.1	12	27	13	5.4	9.2	102	2.2	2.7	1.6	1:1	29	34
	Pu-23	Up	Value		06	34		1.1	6.5	13	3.2	52	117	56	24	39	440	9.2	11	7.0	3.8	126	144
	Ci/g)	HIGH	LOW	1.29	1.01	1.41	1.25	1.50	1.08	1.33	1.50	1.04	1.30	1.63	1.03	1.05	1.30	1.17	1.57	1.49	1.00	1.13	1.17
	240 Activity Concentration (pCi/g)	Down	Uncer.	6.5	18	10	18	1.1	1.6	0.5	0.5	10	30	6.7	4.9	8.1	29	1.1	2.7	9.2	0.5	18	89
Aliquot #3	vity Conce	Do	Value	28	77	43	77	4.9	6.5	2.2	3.2	43	127	41	20	35	126	3.8	12	39	2.7	17	290
		Up	Uncer.	9.8	18	14.0	15	0.5	1.6	0.5	0.5	9.7	23	5.9	4.3	2.6	38	1.1	4.9	5.9	0.5	20	79
,	Pu-239/	ח	Value	36	92	61	62	3.2	7.0	1.6	2.2	42	86	25	19	33	164	3.2	19	27	2.7	87	340
Sampling	Location	-qnS	sample	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sam	Loc	#		31	32	33	34	35	36	37	38	39	40	31	32	33	34	35	36	37	38	39	40
Base	Samile	Number	TAGILITA I	GS0000669	GS0000670	GS0000671	GS0000672	GS0000673	GS0000674	GS0000675	GS0000676	GS0000677	GS0000678	GS0000679	GS0000680	GS0000681	GS0000682	GS0000683	GS0000684	GS0000685	9890000SD	GS0000687	GS0000688

Table A-2. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (2000 Sampling)

	Ι	Т		T	T	Τ-	Т	Т	_	_	Т	Τ	Т	Т		T		т	1		Τ	1	
	Ci/g)	НІСН	TOW						1.15							1.27	1.31	1.29	1.08	1.20	1.17	1.19	2.30
	intration (1	wn	Uncer.						2.2							9.7	36	6.5	1.6	0.5	1.1	31	58
Aliquot #6	vity Conce	Down	Value						8.1							43	154	29	6.5	2.7	3.8	133	251
,	Pu-239/240 Activity Concentration (pCi/g)	c	Uncer.						1.6							9.7	46	5.4	1.6	0.5	1.1	37	133
	Pu-23	$\Omega_{\mathbf{p}}$	Value						7.0							34	201	22	7.0	3.2	3.2	159	575
	Ci/g)	HIGH	TOW						2.00	1.00	2.36	1.10	1.20	1.20	1.26	1.05	1.63	1.13	1.07	2.00	1.33	1.03	1.02
	240 Activity Concentration (pCi/g)	wn	Uncer.						3.2	1.1	3.2	16	21	5.4	3.8	17	52	2.2	2.2	1.1	0.5	21	29
Aliquot #5	vity Conce	Down	Value						14	3.2	14	19	88	23	17	71	226	9.8	8.1	4.3	2.2	88	123
1		d	Uncer.						1.6	1.1	1.6	17	17	4.3	4.9	17	32	2.2	2.2	0.5	0.4	20	28
	Pu-239/	Up	Value						7.0	3.2	5.9	74	73	19	21	75	139	6.7	9.8	2.2	1.6	85	121
Sampling	Location	-qnS	sample	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sam	Loca	#	‡	31	32	33	34	35	36	37	38	39	40	31	32	33	34	35	36	37	38	39	40
, C	Base	Sample	Number	GS0000669	GS0000670	GS0000671	GS0000672	GS0000673	GS0000674	GS0000675	GS0000676	GS0000677	GS0000678	GS0000679	GS0000680	GS0000681	GS0000682	GS0000683	GS0000684	GS0000685	GS0000686	GS0000687	GS0000688

Table A-2. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (2000 Sampling)

	Ci/g)	нісн	MOI																	1.00		1.02	5.40
	Pu-239/240 Activity Concentration (pCi/g)	Λη	Uncer.																	1.6		28	22
Aliquot #8	vity Conce	Down	Value																	5.9		119	92
	9/240 Acti	a	Uncer.																	1.6		27	4.0
	Pu-23	Up	Value																	5.9		117	17
	Ci/g)	HIGH	LOW															1.13	1.56	1.00	1.00	1.05	1.48
	240 Activity Concentration (pCi/g)	Down	Uncer.															1.1	3.2	0.5	1.1	17	52
Aliquot #7	vity Conce	Do	Value															4.9	15	2.2	3.2	74	223
		Up	Uncer.															1:1	2.2	0.5	0.5	16	77
	Pu-239/	n	Value															4.3	6.7	2.2	3.2	70	331
Sampling	Location	-qnS	sample	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sam	Loci	#	11	31	32	33	34	35	36	37	38	39	40	31	32	33	34	35	36	37	38	39	40
Rose	Sample	Number	DOUINAT	GS0000669	GS0000670	GS0000671	GS0000672	GS0000673	GS0000674	GS0000675	GS0000676	GS0000677	GS0000678	GS0000679	GS0000680	GS0000681	GS0000682	GS0000683	GS0000684	GS0000685	GS0000686	GS0000687	GS0000688

Table A-2. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (2000 Sampling)

Aliquot #10	Pu-239/240 Activity Concentration (pCi/g)	Down	Uncer. Value Uncer. LOW																				
	Pu-239	Up	Value																				0.0
	pCi/g)	HIGH	LOW																				01.0
	Pu-239/240 Activity Concentration (pCi/g)	Down	Uncer.																				707
Aliquot #9	ivity Conc	oQ	Value																				0000
	9/240 Acti	$U_{\mathbf{p}}$	Uncer.																				070
	Pu-23	n	Value																				
Sampling	Location	-qnS	sample	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	
Sam	Loc	#	‡	31	32	33	34	35	36	37	38	39	40	31	32	33	34	35	36	37	38	39	100
ţ	Base	Sample	Indilloer	6990000SD	0290000SD	GS0000671	GS0000672	GS0000673	GS0000674	GS0000675	GS0000676	GS0000677	GS0000678	GS0000679	GS0000680	GS0000681	GS0000682	GS0000683	GS0000684	GS0000685	GS0000686	GS0000687	

Table A-2. Individual Aliquot 239/240 Pu Activity Concentrations (Continued) (2000 Sampling)

			,				,		,									-			
Aliquot arameters	Percent Coefficient Of Variation	18	36	34	46	42	51	125	95	19	109	64	23	35	57	42	33	137	25	28	158
Combined Aliquot Statistical Parameters	Mean Activity Concentration (pCi/g)	38	113	59	74	2.8	11	16	4.2	09	237	52	23	48	219	9.5	11	7.5	2.9	104	255
Sampling Location	Sub- sample	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	2-4	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Sam Loca	#	31	32	33	34	35	98	28	88	39	40	31	32	33	34	35	36	37	38	39	40
Base	Number	GS0000669	GS0000670	GS0000671	GS0000672	GS0000673	GS0000674	GS0000675	GS0000676	GS0000677	GS0000078	GS0000679	GS0000680	GS0000681	GS0000682	GS0000683	GS0000684	GS0000685	GS0000686	GS0000687	GS0000688

Blank Cells Represent Samples Where an Aliquot Did Not Exist (Aliquot Number Varied from 2 to 10 per Sample)

Table A-3. University of Pittsburgh - Individual Sub-Aliquot Estimated ^{239/240}Pu Activity Concentrations

Base Sample	Sub-Aliquot	Estimated Pu-239/240 A (pCi/g		Relative Percent			
Number	Number	U. of Pittsburgh**	AFIERA	Difference			
	1	6,979 ± 350					
	2	5,884 + 295					
	3	$3,633 \pm 182$					
	4	5,392 ± 270					
	5	6,963 ± 349					
	6	4,276 ± 214					
GS0000050	7	3,013 ± 151	4,381	0.34			
	8	2,001 ± 100	1,501	0.5 1			
	9	2,417 ± 121					
	10	$3,098 \pm 156$					
	Mean	4,366					
	% CV	42					
	1	2,611 <u>+</u> 131					
	2	$2,055 \pm 103$					
	3	5,279 <u>+</u> 265					
	4	4,878 <u>+</u> 245					
	5	$1,943 \pm 97$					
	6	3,238 ± 163					
GS0000051	7	2,077 ± 104	2,799	2.8			
	8	4,078 ± 204	, , , ,				
	9	$1,223 \pm 62$					
	10	$1,398 \pm 70$					
	Mean	2,878					
	% CV	50					
	1	687 <u>+</u> 131					
	2	585 ± 103					
	3	465 ± 265					
	4	442 ± 245					
	5	503 ± 97					
	6	626 ± 163					
GS0000052	7	747 ± 104	579	- 3.3			
	8	415 ± 204					
	9	872 <u>+</u> 62					
	10	259 <u>+</u> 70					
	Mean	560					
	% CV	32					

Table A-3. University of Pittsburgh - Individual Sub-Aliquot Estimated ^{239/240}Pu Activity Concentrations (Continued)

Base Sample	Sub-Aliquot	Estimated Pu-239/240	Activity Concentration	Relative			
Number	Number	(pCi	(/g)*	Percent			
1 (dillool	ranibei	U. of Pittsburgh**	AFIERA	Difference			
	1	2,655 ± 133					
	2	3,751 <u>+</u> 188					
	3	2,566 <u>+</u> 129					
	4	1,152 ± 58					
	5	2,615 ± 131					
GG0000057	6	3,579 <u>+</u> 179					
GS0000057	7	2,453 ± 123	2,281	5.5			
	8	1,522 ± 76	,				
	9	$2,156 \pm 108$					
	10	1,650 ± 83					
	Mean	2,410					
	% CV	35					
	1	272,436 ± 13,646***					
	2	455 <u>+</u> 23					
	3	447 <u>+</u> 23					
	4	4,321 ± 22					
	5	442 ± 22					
~~~~~	6	534 ± 27					
GS0000079	7	554 <u>+</u> 28	36,123	25			
	8	490 ± 25					
	9	$697 \pm 30$					
	10	342 <u>+</u> 17					
	Mean	28,062					
	% CV	306	,				

^{* 239/240} Pu to 241 Am Ratio of 5.4 Assumed

^{** 95 %} Confidence Level

^{***} Single Particle  $PuO_2$  Estimated Diameter = 250  $\mu m$ 

## Appendix B

Select Plots of ^{239/240}Pu Activity Concentration For Individual Aliquots (This Page Intentionally Left Blank)

Figure B-1. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 67) [Mean  $^{239/240}$ Pu = 271 pCi g⁻¹, % CV = 96%].

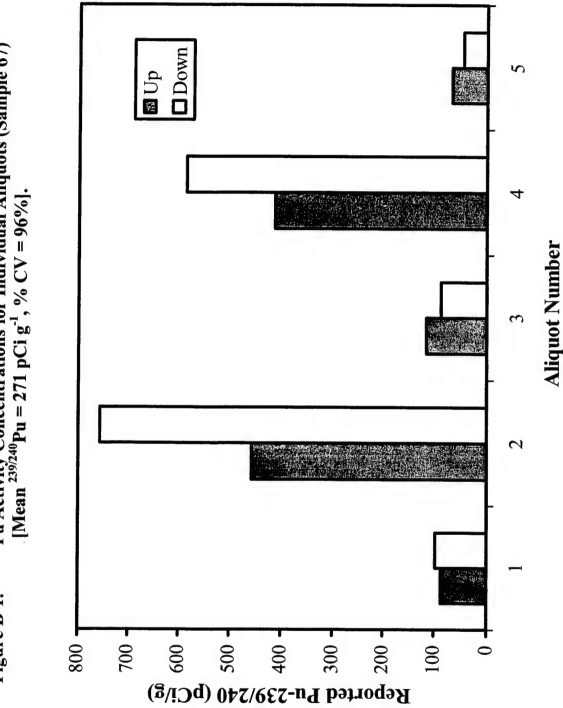


Figure B-2. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 15) [Mean ^{239/240}Pu = 219 pCi g⁻¹, % CV = 128 %].

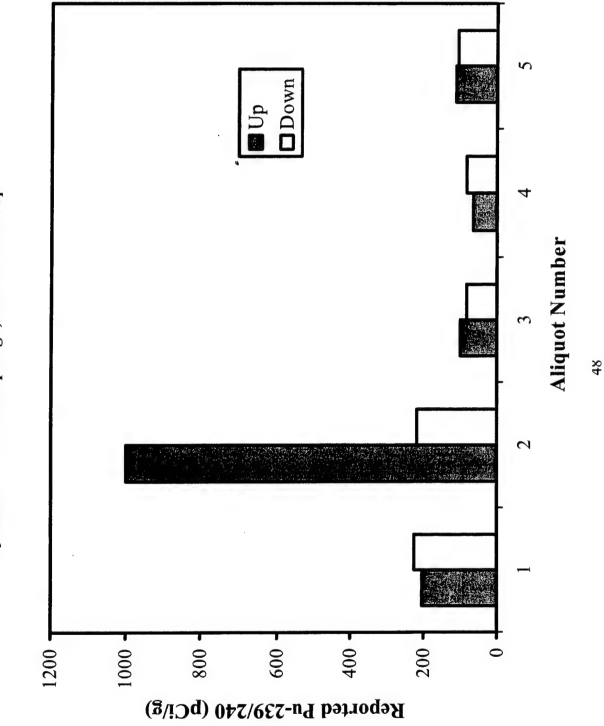


Figure B-3. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 53) [Mean  $^{239/240}$ Pu = 12,900 pCi g⁻¹, % CV = 115 %].

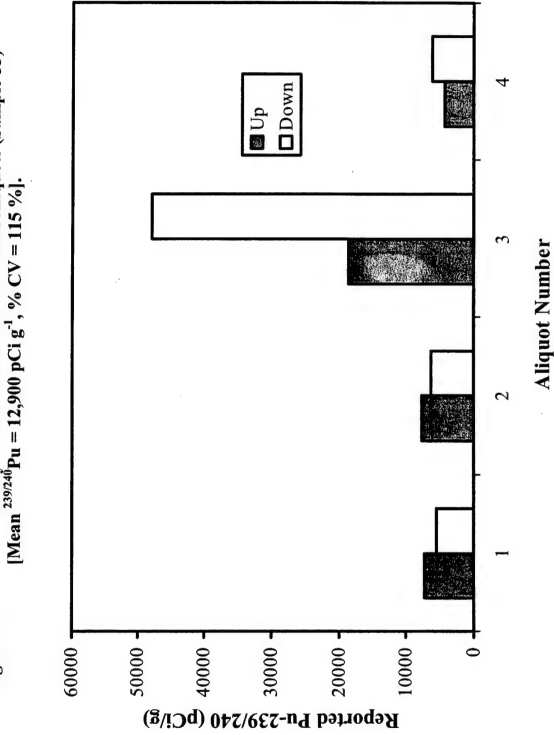


Figure B-4. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 685) [Mean ^{239/240}Pu = 7.5 pCi g⁻¹, % CV = 137 %].

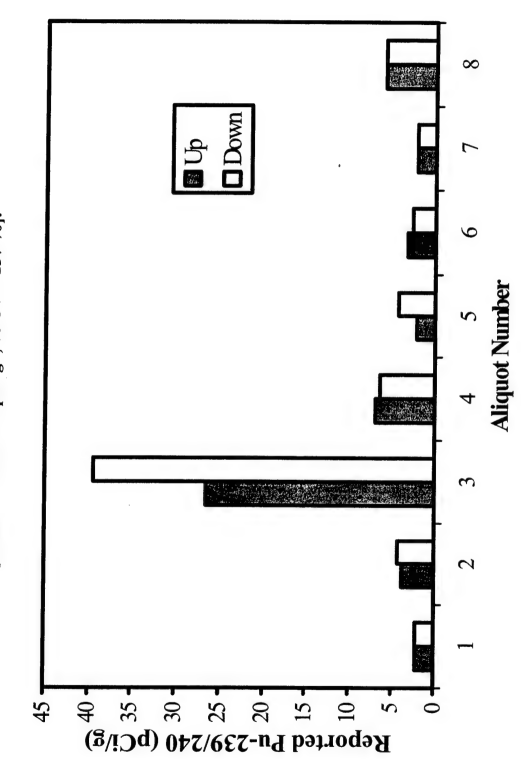


Figure B-5. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 675) [Mean  $^{239/240}$ Pu = 16 pCi  $g^{-1}$ , % CV = 125 %].

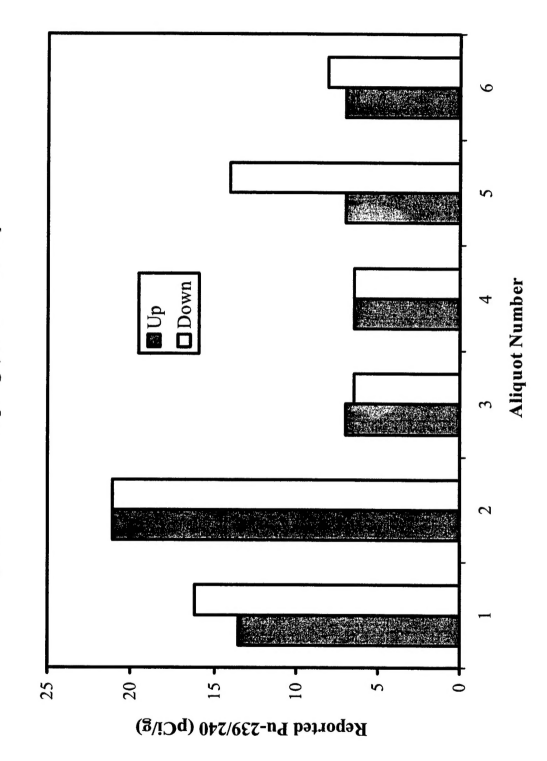


Figure B-6.  $^{239/240}$ Pu Activity Concentrations for Individual Aliquots (Sample 75) [Mean  $^{239/240}$ Pu = 470 pCi g⁻¹, % CV = 72 %].

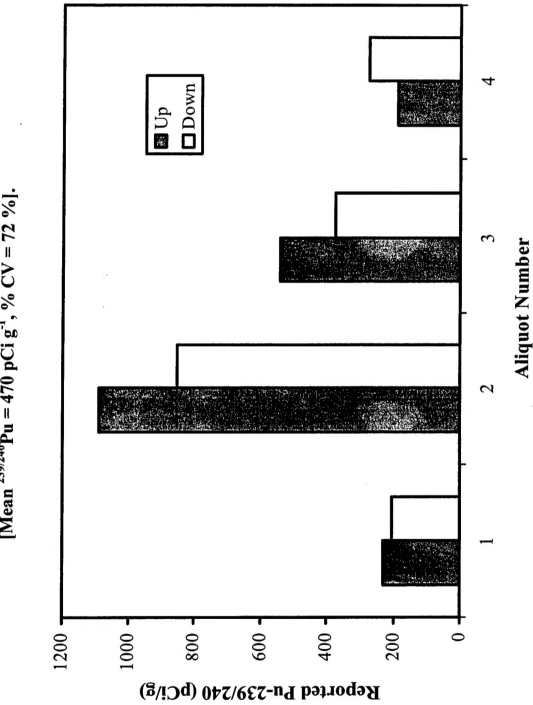


Figure B-7.  $^{239/240}$ Pu Activity Concentrations for Individual Aliquots (Sample 60) [Mean  $^{239/240}$ Pu = 128 pCi g⁻¹, % CV = 13 %].

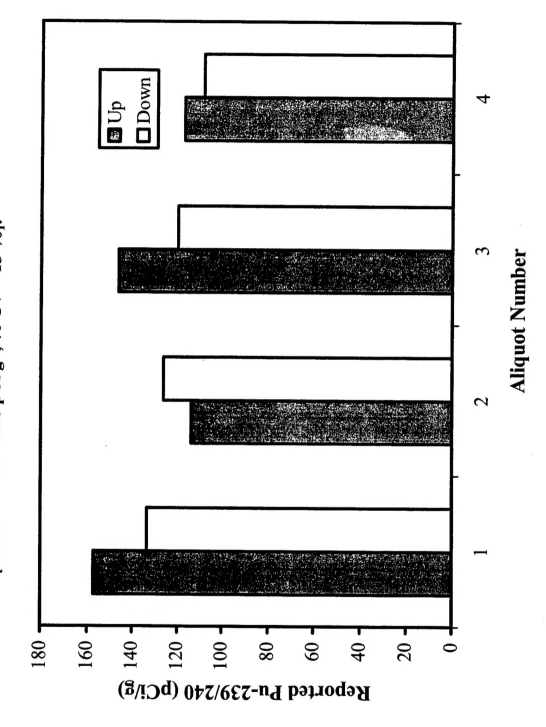


Figure B-8. ^{239/240}Pu Activity Concentrations for Individual Aliquots (Sample 686) [Mean  $^{239/240}$ Pu = 2.9 pCi g⁻¹, % CV = 25 %].

